Biological Product Deviation Codes

Blood BPD Codes or Non-Blood BPD Codes

Blood BPD Codes:

The following list of Biological Product Deviation (BPD) Codes should be used as a tool for assigning a specific code to a reportable event when submitting the report to FDA. This list should not be used solely for the purpose of determining if an event is reportable. You should have a process consistent with the draft guidance document, "Biological Product Deviation Reporting for Blood and Plasma Establishments," to determine which events are reportable. The list includes deviations from regulations, standards, and standard operating procedures (SOPs) that may affect the safety, purity, or potency of a product. The list of codes was developed based on the error and accident reports previously submitted to FDA. These events may not apply to all establishments because they include deviations and unexpected events related to SOPs implemented at individual establishments and may not be an industry standard or a procedure at your facility. The use of BPD Codes will assist the FDA in analyzing the data submitted and streamline the trend analysis process.

Donor Suitability

PD - Post Donation Information

DS - Donor Screening

DD - Donor Deferral

BC - Blood Collection

CP - Component Preparation

Laboratory Testing

VT - Viral Testing

RT - Routine Testing

LA - Labeling

QC - Quality Control and Distribution

MI - Miscellaneous

PD/DS/DD DONOR SUITABILITY

PD--**** POST DONATION INFORMATION

PD-10-** Miscellaneous PD-10-01 Other

PD-11-** Testing

PD-11-01 Other

PD-11-02 Tested positive for Hepatitis B post donation

PD-11-03 Tested positive for Hepatitis B prior to donation

- PD-11-04 Tested positive for Hepatitis C post donation
- **PD-11-05** Tested positive for Hepatitis C prior to donation
- **PD-11-06** Tested positive for HIV post donation
- **PD-11-07** Tested positive for HIV prior to donation
- PD-11-08 Tested positive for HTLV I/II post donation
- **PD-11-09** Tested positive for HTLV I/II prior to donation
- PD-11-10 Tested positive for sexually transmitted disease post donation
- **PD-11-11** Tested positive for sexually transmitted disease prior to donation
- PD-11-12 Tested positive for hepatitis not specified, post donation
- PD-11-13 Tested positive for hepatitis not specified, prior to donation
- PD-11-14 Tested positive at another center, specific testing unknown
- **PD-11-15** Tested positive for Hepatitis A post donation
- **PD-11-16** Tested positive for Hepatitis A prior to donation
- PD-11-17 Elevated ALT post donation
- **PD-11-18** Elevated ALT prior to donation

PD-12-** Behavior/History

- **PD-12-01** Other
- PD-12-02 History of hepatitis not specified
- PD-12-03 History of jaundice
- PD-12-04 History of Hepatitis B
- PD-12-05 History of Hepatitis C
- PD-12-06 Sexually transmitted disease
- PD-12-07 Sex partner has or had a sexually transmitted disease
- **PD-12-08** Sex partner tested positive for HIV
- PD-12-09 Sex partner tested positive for HTLV I/II
- PD-12-10 Sex partner tested positive for HBV
- **PD-12-11** Sex partner tested positive for HCV
- PD-12-12 Sex partner tested positive for hepatitis, not specified
- PD-12-13 Sex partner engaged in high risk behavior
- PD-12-14 Male donor had sex with another man
- PD-12-15 Female had sex with a man who had sex with another man
- **PD-12-16** IV drug use
- PD-12-17 Sex with IV drug user
- PD-12-18 Non-IV-drug use
- PD-12-19 Sex partner used non-IV drugs
- PD-12-20 Donor lived in or immigrated from an HIV Group O risk area
- PD-12-21 Sex partner lived in or immigrated from an HIV Group O risk area
- PD-12-22 Exchanged sex for drugs or money
- PD-12-23 Sex partner exchanged sex for drugs or money
- PD-12-24 Donor received tattoo
- **PD-12-25** Donor received ear piercing
- **PD-12-26** Donor received body piercing
- **PD-12-27** Donor received accidental needlestick
- **PD-12-28** Donor received transfusion or clotting factors
- **PD-12-29** Donor received bone graft or transplant
- **PD-12-30** Donor was exposed to blood or body fluids
- **PD-12-31** Donor had multiple blood or body fluid exposures, e.g., tattoo and piercing
- **PD-12-32** Non-sexual exposure to HIV

- PD-12-33 Non-sexual exposure to hepatitis, type not specified
- **PD-12-34** Non-sexual exposure to Hepatitis B
- **PD-12-35** Non-sexual exposure to Hepatitis C
- PD-12-36 Travel to malaria endemic area/history of malaria
- **PD-12-37** History of disease or surgery
- **PD-12-38** History of cancer
- PD-12-39 History of Creutzfeldt-Jakob Disease
- $\textbf{PD-12-40} \ \textbf{Risk factors associated with Creutzfeldt-Jakob Disease brain surgery}$
- PD-12-41 Risk factors associated with Creutzfeldt-Jakob Disease family history
- PD-12-42 Risk factors associated with Creutzfeldt-Jakob Disease (vCJD) travel
- PD-12-43 Risk factors associated with Creutzfeldt-Jakob Disease received insulin
- **PD-12-44** Received growth hormone
- PD-12-45 Received Proscar, Tegison or Accutane
- PD-12-46 Received medication or antibiotics
- PD-12-47 Received vaccine or immune globulin
- PD-12-48 Exposure to a disease
- PD-12-49 Incarcerated
- **PD-12-50** Resided in a rehabilitation center or psychiatric hospital
- **PD-12-51** History of Hepatitis A
- **PD-12-52** Exposure to Hepatitis A
- PD-12-53 Multiple high risk behaviors/contacts
- PD-12-54 Positive drug screen
- PD-12-55 Deferred by another center

PD-13-** Illness

- **PD-13-01** Post donation illness (not hepatitis, HIV, HTLV-I, STD, cancer or cold/flu related)
- PD-13-02 Post donation diagnosis or symptoms of Hepatitis B
- PD-13-03 Post donation diagnosis or symptoms of Hepatitis C
- PD-13-04 Post donation diagnosis or symptoms of HIV
- PD-13-05 Post donation diagnosis or symptoms of HTLV I/II
- PD-13-06 Post donation diagnosis or symptoms of sexually transmitted disease
- PD-13-07 Post donation diagnosis or symptoms of hepatitis, not specified
- PD-13-08 Post donation diagnosis or symptoms of Hepatitis A
- PD-13-09 Post donation diagnosis of cancer
- **PD-14-**** Not specifically related to high risk behavior or unsuitable history
 - **PD-14-01** Other
 - PD-14-02 Donor does not want their blood used
 - **PD-14-03** Donated to be tested or called back for test results

DS--**** DONOR SCREENING

DS-20-** Miscellaneous

DS-20-01 Other

DS-21-** Donor did not meet acceptance criteria

DS-21-01 Other

DS-21-02 Hemoglobin or Hematocrit unacceptable, not documented, or testing performed incorrectly

DS-21-03 Temperature unacceptable or not documented

DS-21-04 Medical review or physical not performed or inadequate

DS-21-05 Platelet count, no documented platelet count for product

DS-21-06 Unexplained weight loss

DS-22-** Donor record incomplete or incorrect

DS-22-01 Other

DS-22-02 Donor identification

DS-22-03 Donor history questions

DS-22-04 Arm inspection

DS-22-05 Donor signature missing

DS-22-06 Confidential Unit Exclusion (CUE) procedure not performed in accordance with specifications

DS-22-07 Donor confidentiality compromised

DS-23-** Deferral screening not done

DS-23-01 Donor not previously deferred

DS-24-** Deferral screening not done, donor previously deferred due to testing:

DS-24-01 Other

DS-24-02 HIV reactive

DS-24-03 HBsAg reactive

DS-24-04 Anti-HBc reactive

DS-24-05 Anti-HCV reactive

DS-24-06 Anti-HTLV-I reactive

DS-24-07 ALT elevated

DS-24-08 Syphilis reactive

DS-25-** Deferral screening not done, donor previously deferred due to history

DS-25-01 Other

DS-25-02 History of hepatitis, not specified

DS-25-03 History of jaundice

DS-25-04 History of Hepatitis B

DS-25-05 History of Hepatitis C

- DS-25-06 Sexually transmitted disease
- **DS-25-07** Sex partner has or had a sexually transmitted disease
- **DS-25-08** Sex partner tested positive for HIV
- **DS-25-09** Sex partner tested positive for HTLV I/II
- **DS-25-10** Sex partner tested positive for HBV
- **DS-25-11** Sex partner tested positive for HCV
- **DS-25-12** Sex partner tested positive for hepatitis, not specified
- **DS-25-13** Sex partner engaged in high risk behavior
- DS-25-14 Male donor had sex with another man
- **DS-25-15** Female had sex with a man who had sex with another man
- **DS-25-16** IV drug use
- DS-25-17 Sex with IV drug user
- **DS-25-18** Non-IV-drug use
- **DS-25-19** Sex partner used non-IV drugs
- DS-25-20 Donor lived in or immigrated from an HIV Group O risk area
- DS-25-21 Sex partner lived in or immigrated from an HIV Group O risk area
- **DS-25-22** Exchanged sex for drugs or money
- **DS-25-23** Sex partner exchanged sex for drugs or money
- DS-25-24 Donor received tattoo
- **DS-25-25** Donor received ear piercing
- **DS-25-26** Donor received body piercing
- **DS-25-27** Donor received accidental needlestick
- **DS-25-28** Donor received transfusion or clotting factors
- **DS-25-29** Donor received bone graft or transplant
- **DS-25-30** Donor was exposed to blood or body fluids
- **DS-25-31** Donor had multiple blood or body fluid exposures, e.g., tattoo and piercing
- **DS-25-32** Non-sexual exposure to HIV
- DS-25-33 Non-sexual exposure to hepatitis, type not specified
- **DS-25-34** Non-sexual exposure to Hepatitis B
- **DS-25-35** Non-sexual exposure to Hepatitis C
- **DS-25-36** Travel to malaria endemic area/history of malaria
- **DS-25-37** History of disease or surgery
- DS-25-38 History of cancer
- **DS-25-39** History of Creutzfeldt-Jakob Disease
- **DS-25-40** Risk factors associated with Creutzfeldt-Jakob Disease brain surgery
- **DS-25-41** Risk factors associated with Creutzfeldt-Jakob Disease family history
- **DS-25-42** Risk factors associated with Creutzfeldt-Jakob Disease (vCJD) travel
- **DS-25-43** Risk factors associated with Creutzfeldt-Jakob Disease received insulin
- **DS-25-44** Received growth hormone
- **DS-25-45** Received Proscar, Tegison or Accutane
- DS-25-46 Received medication or antibiotics
- **DS-25-47** Received vaccine or immune globulin
- **DS-25-48** Exposure to a disease
- **DS-25-49** Incarcerated
- **DS-25-50** Resided in a rehabilitation center or psychiatric hospital
- **DS-25-51** History of Hepatitis A
- **DS-25-52** Exposure to Hepatitis A
- **DS-25-53** Multiple high risk behaviors/contacts

DS-25-54 Positive drug screen

DS-25-55 Deferred by another center

DS-26-** Incorrect ID used during deferral search

DS-26-01 donor not previously deferred

DS-27-** Incorrect ID used during deferral search, donor previously deferred due to testing

DS-27-01 Other

DS-27-02 HIV reactive

DS-27-03 HBsAg reactive

DS-27-04 Anti-HBc reactive

DS-27-05 Anti-HCV reactive

DS-27-06 Anti-HTLV-I reactive

DS-27-07 ALT elevated

DS-27-08 Syphilis reactive

DS-28-** Incorrect ID used during deferral search, donor previously deferred due to history

DS-28-01 Other

DS-28-02 History of hepatitis, not specified

DS-28-03 History of jaundice

DS-28-04 History of Hepatitis B

DS-28-05 History of Hepatitis C

DS-28-06 Sexually transmitted disease

DS-28-07 Sex partner has or had a sexually transmitted disease

DS-28-08 Sex partner tested positive for HIV

DS-28-09 Sex partner tested positive for HTLV I/II

DS-28-10 Sex partner tested positive for HBV

DS-28-11 Sex partner tested positive for HCV

DS-28-12 Sex partner tested positive for hepatitis, not specified

DS-28-13 Sex partner engaged in high risk behavior

DS-28-14 Male donor had sex with another man

DS-28-15 Female had sex with a man who had sex with another man

DS-28-16 IV drug use

DS-28-17 Sex with IV drug user

DS-28-18 Non-IV-drug use

DS-28-19 Sex partner used non-IV drugs

DS-28-20 Donor lived in or immigrated from an HIV Group O risk area

DS-28-21 Sex partner lived in or immigrated from an HIV Group O risk area

DS-28-22 Exchanged sex for drugs or money

DS-28-23 Sex partner exchanged sex for drugs or money

DS-28-24 Donor received tattoo

DS-28-25 Donor received ear piercing

DS-28-26 Donor received body piercing

DS-28-27 Donor received accidental needlestick

DS-28-28 Donor received transfusion or clotting factors

- **DS-28-29** Donor received bone graft or transplant
- **DS-28-30** Donor was exposed to blood or body fluids
- **DS-28-31** Donor had multiple blood or body fluid exposures, e.g., tattoo and piercing
- **DS-28-32** Non-sexual exposure to HIV
- **DS-28-33** Non-sexual exposure to hepatitis, type not specified
- **DS-28-34** Non-sexual exposure to Hepatitis B
- **DS-28-35** Non-sexual exposure to Hepatitis C
- **DS-28-36** Travel to malaria endemic area/history of malaria
- **DS-28-37** History of disease or surgery
- **DS-28-38** History of cancer
- DS-28-39 History of Creutzfeldt-Jakob Disease
- **DS-28-40** Risk factors associated with Creutzfeldt-Jakob Disease brain surgery
- **DS-28-41** Risk factors associated with Creutzfeldt-Jakob Disease family history
- DS-28-42 Risk factors associated with Creutzfeldt-Jakob Disease (vCJD) travel
- **DS-28-43** Risk factors associated with Creutzfeldt-Jakob Disease received insulin
- **DS-28-44** Received growth hormone
- DS-28-45 Received Proscar, Tegison or Accutane
- **DS-28-46** Received medication or antibiotics
- DS-28-47 Received vaccine or immune globulin
- **DS-28-48** Exposure to a disease
- **DS-28-49** Incarcerated
- DS-28-50 Resided in a rehabilitation center or psychiatric hospital
- **DS-28-51** History of Hepatitis A
- **DS-28-52** Exposure to Hepatitis A
- **DS-28-53** Multiple high risk behaviors/contacts
- **DS-28-54** Positive drug screen
- **DS-28-55** Deferred by another center

DS-29-** Donor gave history which warranted deferral and was not deferred

- **DS-29-01** Other
- DS-29-02 History of hepatitis, not specified
- **DS-29-03** History of jaundice
- **DS-29-04** History of Hepatitis B
- **DS-29-05** History of Hepatitis C
- **DS-29-06** Sexually transmitted disease
- DS-29-07 Sex partner has or had a sexually transmitted disease
- **DS-29-08** Sex partner tested positive for HIV
- **DS-29-09** Sex partner tested positive for HTLV I/II
- **DS-29-10** Sex partner tested positive for HBV
- DS-29-11 Sex partner tested positive for HCV
- **DS-29-12** Sex partner tested positive for hepatitis, not specified
- **DS-29-13** Sex partner engaged in high risk behavior
- **DS-29-14** Male donor had sex with another man
- DS-29-15 Female had sex with a man who had sex with another man
- **DS-29-16** IV drug use
- DS-29-17 Sex with IV drug user
- DS-29-18 Non-IV-drug use
- **DS-29-19** Sex partner used non-IV drugs

DS-29-20 Donor lived in or immigrated from an HIV Group O risk area

DS-29-21 Sex partner lived in or immigrated from an HIV Group O risk area

DS-29-22 Exchanged sex for drugs or money

DS-29-23 Sex partner exchanged sex for drugs or money

DS-29-24 Donor received tattoo

DS-29-25 Donor received ear piercing

DS-29-26 Donor received body piercing

DS-29-27 Donor received accidental needlestick

DS-29-28 Donor received transfusion or clotting factors

DS-29-29 Donor received bone graft or transplant

DS-29-30 Donor was exposed to blood or body fluids

DS-29-31 Donor had multiple blood or body fluid exposures, e.g., tattoo and piercing

DS-29-32 Non-sexual exposure to HIV

DS-29-33 Non-sexual exposure to hepatitis, type not specified

DS-29-34 Non-sexual exposure to Hepatitis B

DS-29-35 Non-sexual exposure to Hepatitis C

DS-29-36 Travel to malaria endemic area/history of malaria

DS-29-37 History of disease or surgery

DS-29-38 History of cancer

DS-29-39 History of Creutzfeldt-Jakob Disease

DS-29-40 Risk factors associated with Creutzfeldt-Jakob Disease - brain surgery

DS-29-41 Risk factors associated with Creutzfeldt-Jakob Disease - family history

DS-29-42 Risk factors associated with Creutzfeldt-Jakob Disease (vCJD) - travel

DS-29-43 Risk factors associated with Creutzfeldt-Jakob Disease - received insulin

DS-29-44 Received growth hormone

DS-29-45 Received Proscar, Tegison or Accutane

DS-29-46 Received medication or antibiotics

DS-29-47 Received vaccine or immune globulin

DS-29-48 Exposure to a disease

DS-29-49 Incarcerated

DS-29-50 Resided in a rehabilitation center or psychiatric hospital

DS-29-51 History of Hepatitis A

DS-29-52 Exposure to Hepatitis A

DS-29-53 Multiple high risk behaviors/contacts

DS-29-54 Positive drug screen

DS-29-55 Deferred by another center

DD-**-** DONOR DEFERRAL

DD-30-** Miscellaneous

DD-30-01 Other

DD-31-** Donor missing or incorrectly identified on deferral list, donor was or should have been previously deferred due to testing:

DD-31-01 Other

DD-31-02 HIV reactive

DD-31-03 HBsAg reactive

DD-31-04 Anti-HBc reactive

DD-31-05 Anti-HCV reactive

DD-31-06 Anti-HTLV-I reactive

DD-31-07 ALT elevated

DD-31-08 Syphilis reactive

DD-32-** Donor missing or incorrectly identified on deferral list, donor was or should have been previously deferred due to history:

DD-32-01 Other

DD-32-02 History of hepatitis, not specified

DD-32-03 History of jaundice

DD-32-04 History of Hepatitis B

DD-32-05 History of Hepatitis C

DD-32-06 Sexually transmitted disease

DD-32-07 Sex partner has or had sexually transmitted disease

DD-32-08 Sex partner tested positive for HIV

DD-32-09 Sex partner tested positive for HTLV I/II

DD-32-10 Sex partner tested positive for HBV

DD-32-11 Sex partner tested positive for HCV

DD-32-12 Sex partner tested positive for hepatitis, not specified

DD-32-13 Sex partner engaged in high risk behavior

DD-32-14 Male donor had sex with another man

DD-32-15 Female had sex with a man who had sex with another man

DD-32-16 IV drug use

DD-32-17 Sex with IV drug user

DD-32-18 Non-IV-drug use

DD-32-19 Sex partner used non-IV drugs

DD-32-20 Donor lived in or immigrated from an HIV Group O risk area

DD-32-21 Sex partner lived in or immigrated from an HIV Group O risk area

DD-32-22 Exchanged sex for drugs or money

DD-32-23 Sex partner exchanged sex for drugs or money

DD-32-24 Donor received tattoo

DD-32-25 Donor received ear piercing

DD-32-26 Donor received body piercing

DD-32-27 Donor received accidental needlestick

DD-32-28 Donor received transfusion or clotting factors

DD-32-29 Donor received bone graft or transplant

DD-32-30 Donor was exposed to blood or body fluids

DD-32-31 Donor had multiple blood or body fluid exposures, e.g., tattoo and piercing

DD-32-32 Non-sexual exposure to HIV

DD-32-33 Non-sexual exposure to hepatitis, type not specified

DD-32-34 Non-sexual exposure to Hepatitis B

DD-32-35 Non-sexual exposure to Hepatitis C

DD-32-36 Travel to malaria endemic area/history of malaria

DD-32-37 History of disease or surgery

DD-32-38 History of cancer

DD-32-39 History of Creutzfeldt-Jakob Disease

DD-32-40 Risk factors associated with Creutzfeldt-Jakob Disease - brain surgery

DD-32-41 Risk factors associated with Creutzfeldt-Jakob Disease - family history

DD-32-42 Risk factors associated with Creutzfeldt-Jakob Disease (vCJD) - travel

DD-32-43 Risk factors associated with Creutzfeldt-Jakob Disease - received insulin

DD-32-44 Received growth hormone

DD-32-45 Received Proscar, Tegison or Accutane

DD-32-46 Received medication or antibiotics

DD-32-47 Received vaccine or immune globulin

DD-32-48 Exposure to a disease

DD-32-49 Incarcerated

DD-32-50 Resided in a rehabilitation center or psychiatric hospital

DD-32-51 History of Hepatitis A

DD-32-52 Exposure to Hepatitis A

DD-32-53 Multiple high risk behaviors/contacts

DD-32-54 Positive drug screen

DD-32-55 Deferred by another center

DD-34-** Donor incorrectly deleted from deferral list or donor not reentered properly, donor previously deferred due to testing:

DD-34-01 Other

DD-34-02 HIV reactive

DD-34-03 HBsAg reactive

DD-34-04 Anti-HBc reactive

DD-34-05 Anti-HCV reactive

DD-34-06 Anti-HTLV-I reactive

DD-34-07 ALT elevated

DD-34-08 Syphilis reactive

DD-35-** Donor incorrectly deleted from deferral list, donor previously deferred due to history:

DD-35-01 Other

DD-35-02 History of hepatitis, not specified

DD-35-03 History of jaundice

DD-35-04 History of Hepatitis B

DD-35-05 History of Hepatitis C

DD-35-06 Sexually transmitted disease

DD-35-07 Sex partner has or had a sexually transmitted disease

DD-35-08 Sex partner tested positive for HIV

DD-35-09 Sex partner tested positive for HTLV I/II

DD-35-10 Sex partner tested positive for HBV

DD-35-11 Sex partner tested positive for HCV

DD-35-12 Sex partner tested positive for hepatitis, not specified

DD-35-13 Sex partner engaged in high risk behavior

DD-35-14 Male donor had sex with another man

- **DD-35-15** Female had sex with a man who had sex with another man
- **DD-35-16** IV drug use
- **DD-35-17** Sex with IV drug user
- DD-35-18 Non-IV-drug use
- **DD-35-19** Sex partner used non-IV drugs
- **DD-35-20** Donor lived in or immigrated from an HIV Group O risk area
- **DD-35-21** Sex partner lived in or immigrated from an HIV Group O risk area
- **DD-35-22** Exchanged sex for drugs or money
- DD-35-23 Sex partner exchanged sex for drugs or money
- **DD-35-24** Donor received tattoo
- DD-35-25 Donor received ear piercing
- **DD-35-26** Donor received body piercing
- **DD-35-27** Donor received accidental needlestick
- **DD-35-28** Donor received transfusion or clotting factors
- **DD-35-29** Donor received bone graft or transplant
- **DD-35-30** Donor was exposed to blood or body fluids
- **DD-35-31** Donor had multiple blood or body fluid exposures, e.g., tattoo and piercing
- **DD-35-32** Non-sexual exposure to HIV
- **DD-35-33** Non-sexual exposure to hepatitis, type not specified
- **DD-35-34** Non-sexual exposure to Hepatitis B
- DD-35-35 Non-sexual exposure to Hepatitis C
- **DD-35-36** Travel to malaria endemic area/history of malaria
- DD-35-37 History of disease or surgery
- **DD-35-38** History of cancer
- DD-35-39 History of Creutzfeldt-Jakob Disease
- **DD-35-40** Risk factors associated with Creutzfeldt-Jakob Disease brain surgery
- **DD-35-41** Risk factors associated with Creutzfeldt-Jakob Disease family history
- DD-35-42 Risk factors associated with Creutzfeldt-Jakob Disease (vCJD) travel
- **DD-35-43** Risk factors associated with Creutzfeldt-Jakob Disease received insulin
- DD-35-44 Received growth hormone
- DD-35-45 Received Proscar, Tegison or Accutane
- **DD-35-46** Received medication or antibiotics
- **DD-35-47** Received vaccine or immune globulin
- **DD-35-48** Exposure to a disease
- DD-35-49 Incarcerated
- **DD-35-50** Resided in a rehabilitation center or psychiatric hospital
- **DD-35-51** History of Hepatitis A
- **DD-35-52** Exposure to Hepatitis A
- **DD-35-53** Multiple high risk behaviors/contacts
- **DD-35-54** Positive drug screen
- **DD-35-55** Deferred by another center

BC--**** BLOOD COLLECTION

BC-40-** Miscellaneous

BC-40-01 Other

BC-41-** Sterility compromised

BC-41-01 Other

BC-41-02 Bacterial contamination (identify organism if possible)

BC-41-03 Air contamination

BC-41-04 Arm prep not performed or performed inappropriately

BC-42-** Collection bag

BC-42-01 Other

BC-42-02 Blood drawn into outdated bag

BC-42-03 Incorrect anticoagulant

BC-42-04 Outdated anticoagulant

BC-42-05 Potential collection set (bag, tubing) defect (e.g., product leaking)

BC-42-06 Incorrect collection bag used (e.g., 500 ml bag used instead of 450 ml bag)

BC-43-** Collection process

BC-43-01 Other

BC-43-02 Collection time extended, discrepant, or not documented; not discovered prior to component preparation

BC-43-03 Overbleed; not discovered prior to component preparation

BC-43-04 Collection status not documented or discrepant

BC-43-05 Product contained clots, not discovered prior to distribution

BC-43-06 Product hemolyzed, not discovered prior to distribution

BC-43-07 Source Plasma from two different donors pooled into one pooling bottle

BC-44-** Apheresis collection device

BC-44-01 Other

BC-44-02 Device defect

BC-44-03 Softgoods defect (bags, tubing, etc)

CP--**** COMPONENT PREPARATION

CP-50-** Miscellaneous

CP-50-01 Other

CP-51-** Sterility compromised

CP-51-01 Other

CP-51-02 Bacterial contamination (identify organism if possible)

CP-51-03 Air contamination

CP-51-04 Product integrity compromised during component preparation (e.g., leaking at sterile connection site)

CP-52-** Component not prepared in accordance with specifications

CP-52-01 Other

CP-52-02 Platelets made from Whole Blood collected from donor who took medication that may affect platelet function

CP-52-03 Resting time requirements not met for Platelets

CP-52-04 Platelets not agitated

CP-52-05 Platelet count or platelet yield not acceptable or platelet count not performed on Platelet product

CP-52-06 Processed at incorrect centrifuge setting

CP-52-07 Product not frozen within the appropriate time frame or freezing time not documented

CP-52-08 Product prepared at incorrect temperature or held at incorrect temperature during component preparation

CP-52-09 Washing/deglycerolization not performed in accordance with specifications

CP-52-10 Leukoreduction not performed in accordance with specifications

CP-52-11 Irradiation not performed in accordance with specifications

CP-52-12 Components not prepared within appropriate time frame after collection

CP-52-13 Additive solution not added, added incorrectly, or added to incorrect product

CP-52-14 Thawing frozen product not performed in accordance with specifications

CP-52-15 Pooling not performed in accordance with specifications

CP-52-16 Aliquot preparation not performed in accordance with specifications

CP-53-** Component prepared from Whole Blood unit that was

CP-53-01 Other

CP-53-02 Overweight

CP-53-03 Underweight

CP-53-04 Collected or stored at unacceptable or undocumented temperature

CP-53-05 A difficult collection or had an extended collection time

CP-54-** Component manufactured that was

CP-54-01 Other

CP-54-02 Overweight

CP-54-03 Underweight

VT/RT LABORATORY TESTING

VT-**-** VIRAL TESTING

VT-70-** Miscellaneous

VT-70-01 Other

VT-71-** Testing performed incorrectly for:

VT-71-01 HBsAg

VT-71-02 Anti-HIV-1

VT-71-03 Anti-HIV-2

VT-71-04 Anti-HIV-1/2

VT-71-05 HIV Antigen

VT-71-06 Syphilis

VT-71-07 Anti-HTLV-I/II

VT-71-08 Anti-HBc

VT-71-09 ALT

VT-71-10 Anti-HCV

VT-71-11 More than 1 test, e.g., all viral markers

VT-71-12 Cytomegalovirus

VT-71-13 HIV Nucleic Acid Test (NAT)

VT-71-14 HCV Nucleic Acid Test (NAT)

VT-71-15 HIV/HCV Nucleic Acid Test (NAT)

VT-72-** Sample identification

VT-72-01 Other

VT-72-02 Incorrect sample tested

VT-72-03 Sample used for testing was incorrectly or incompletely labeled

VT-72-04 Unsuitable sample used for testing

RT--**** ROUTINE TESTING

RT-60-** Miscellaneous

RT-60-01 Other

RT-61-** Testing performed incorrectly for:

RT-61-01 Other

RT-61-02 ABO

RT-61-03 Rh

RT-61-04 ABO & Rh

RT-61-05 Antibody screening or identification

RT-61-06 Antigen typing

RT-61-07 Platelet count

RT-61-08 Compatibility

RT-61-09 ABO, Rh, and antibody screen

RT-61-10 ABO, Rh, antibody screen, and compatibility

RT-61-11 Antibody screen and compatibility

RT-62-** Sample identification

RT-62-01 Other

RT-62-02 Incorrect sample tested

RT-62-03 Sample used for testing was incorrectly or incompletely labeled

RT-62-04 Unsuitable sample used for testing (e.g., too old)

RT-63-** Testing performed using reagents in which QC was unacceptable or expired reagents were used

RT-63-01 Other

RT-63-02 ABO

RT-63-03 Rh

RT-63-04 ABO & Rh

RT-63-05 Antibody screening or identification

RT-63-06 Antigen typing

RT-63-07 Multiple testing

LA-**-** LABELING

LA-80-** Miscellaneous

LA-80-01 Other

LA-81-** Labels applied to blood unit incorrect or missing information

LA-81-01 Other

LA-81-02 ABO and/or Rh incorrect

LA-81-03 ABO and/or Rh missing

LA-81-04 Product type incorrect (e.g., RBC labeled as Whole Blood)

LA-81-05 Product type missing

LA-81-06 Extended expiration date or time

LA-81-07 Missing expiration date or time

LA-81-08 Anticoagulant incorrect or missing

LA-81-09 Donor number incorrect or missing

LA-81-10 Multiple labels incorrect or missing

LA-81-11 Volume or weight incorrect or missing

LA-81-12 Irradiation status incorrect or missing

LA-81-13 Leukoreduction status incorrect or missing

LA-81-14 Irradiation and leukoreduction status incorrect or missing

LA-82-** Crossmatch tag or tie tag labels incorrect or missing information

LA-82-01 Other

LA-82-02 Unit ABO and/or Rh incorrect or missing

LA-82-03 Recipient ABO and/or Rh incorrect or missing

LA-82-04 Product type incorrect or missing

LA-82-05 Expiration date or time extended or missing

LA-82-06 Unit or pool number incorrect or missing

LA-82-07 Recipient identification incorrect or missing (specify if autologous unit)

LA-82-08 Antigen incorrect or missing

LA-82-09 Antibody incorrect or missing

LA-82-10 Platelet count incorrect or missing

LA-82-11 HLA type incorrect or missing

LA-82-12 Volume or weight incorrect or missing

LA-82-13 CMV status incorrect or missing

LA-82-14 Irradiation status incorrect or missing

LA-82-15 Leukoreduced status incorrect or missing

LA-82-16 Crossmatch tag switched, both units intended for the same patient

LA-82-17 Crossmatch tag incorrect or missing

LA-82-18 Biohazard or test status incorrect or missing

LA-82-19 Multiple labels incorrect or missing

LA-83-** Transfusion record (crossmatch slip) incorrect or missing information

LA-83-01 Other

LA-83-02 Unit ABO and/or Rh incorrect or missing

LA-83-03 Recipient ABO and/or Rh incorrect or missing

LA-83-04 Product type incorrect or missing

LA-83-05 Expiration date or time extended or missing

LA-83-06 Unit or pool number incorrect or missing

LA-83-07 Recipient identification incorrect or missing (specify if autologous unit)

LA-83-08 Antigen incorrect or missing

LA-83-09 Antibody incorrect or missing

LA-83-10 Platelet count incorrect or missing

LA-83-11 HLA type incorrect or missing

LA-83-12 Volume or weight incorrect or missing

LA-83-13 CMV status incorrect or missing

LA-83-14 Irradiation status incorrect or missing

LA-83-15 Leukoreduced status incorrect or missing

LA-83-16 Transfusion record switched, both units intended for the same patient

LA-83-17 Incorrect transfusion record released with unit (e.g., intended for different patient)

LA-83-18 Biohazard or test status incorrect or missing

LA-83-19 Multiple labels incorrect or missing

QC-**-** QUALITY CONTROL and DISTRIBUTION

QC-90-** Miscellaneous

QC-90-01 Other

QC-91-** Failure to quarantine unit due to medical history:

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QC-91-01 Other
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QC-91-02 History of hepatitis, not specified

QC-91-03 History of jaundice

QC-91-04 History of Hepatitis B

QC-91-05 History of Hepatitis C

QC-91-06 Sexually transmitted disease

QC-91-07 Sex partner has or had a sexually transmitted disease

QC-91-08 Sex partner tested positive for HIV

QC-91-09 Sex partner tested positive for HTLV I/II

QC-91-10 Sex partner tested positive for HBV

QC-91-11 Sex partner tested positive for HCV

QC-91-12 Sex partner tested positive for hepatitis, not specified

QC-91-13 Sex partner engaged in high risk behavior

QC-91-14 Male donor had sex with another man

QC-91-15 Female had sex with a man who had sex with another man

QC-91-16 IV drug use

QC-91-17 Sex with IV drug user

QC-91-18 Non-IV-drug use

QC-91-19 Sex partner used non-IV drugs

QC-91-20 Donor lived in or immigrated from an HIV Group O risk area

QC-91-21 Sex partner lived in or immigrated from an HIV Group O risk area

QC-91-22 Exchanged sex for drugs or money

QC-91-23 Sex partner exchanged sex for drugs or money

QC-91-24 Donor received tattoo

QC-91-25 Donor received ear piercing

QC-91-26 Donor received body piercing

QC-91-27 Donor received accidental needlestick

QC-91-28 Donor received transfusion or clotting factors

QC-91-29 Donor received bone graft or transplant

QC-91-30 Donor was exposed to blood or body fluids

QC-91-31 Donor had multiple blood or body fluid exposures, e.g., tattoo and piercing

QC-91-32 Non-sexual exposure to HIV

OC-91-33 Non-sexual exposure to hepatitis, type not specified

QC-91-34 Non-sexual exposure to Hepatitis B

QC-91-35 Non-sexual exposure to Hepatitis C

QC-91-36 Travel to malaria endemic area/history of malaria

QC-91-37 History of disease or surgery

QC-91-38 History of cancer

QC-91-39 History of Creutzfeldt-Jakob Disease

QC-91-40 Risk factors associated with Creutzfeldt-Jakob Disease - brain surgery

QC-91-41 Risk factors associated with Creutzfeldt-Jakob Disease - family history

QC-91-42 Risk factors associated with Creutzfeldt-Jakob Disease (vCJD) - travel

QC-91-43 Risk factors associated with Creutzfeldt-Jakob Disease - received insulin

QC-91-44 Received growth hormone

OC-91-45 Received Proscar, Tegison or Accutane

QC-91-46 Received medication or antibiotics

QC-91-47 Received vaccine or immune globulin

OC-91-48 Exposure to a disease

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QC-91-50 Resided in a rehabilitation center or psychiatric hospital
       QC-91-51 History of Hepatitis A
       QC-91-52 Exposure to Hepatitis A
       QC-91-53 Multiple high risk behaviors/contacts
       QC-91-54 Positive drug screen
       QC-91-55 Deferred by another center
       QC-91-56 Post donation illness
QC-92-**Required testing incomplete or positive for:
       QC-92-01 Other
       QC-92-02 HIV
       QC-92-03 HBsAg
       QC-92-04 Anti-HBc
       QC-92-05 Anti-HCV
       QC-92-06 Anti-HTLV-I
       QC-92-07 ALT
       QC-92-08 ABO (donor/unit or recipient)
       QC-92-09 Rh (donor/unit or recipient)
       QC-92-10 Antibody screen or identification (donor/unit or recipient)
       QC-92-11 Antigen screen
       QC-92-12 Syphilis
       QC-92-13 All viral markers
       QC-92-14 Compatibility
       QC-92-15 HIV/HCV Nucleic Acid Test (NAT)
QC-93-** Required testing not performed or documented for:
       QC-93-01 Other
       QC-93-02 HIV
       QC-93-03 HBsAg
       QC-93-04 Anti-HBc
       QC-93-05 Anti-HCV
       QC-93-06 Anti-HTLV-I
       QC-93-07 ALT
       QC-93-08 ABO (donor/unit or recipient)
       QC-93-09 Rh (donor/unit or recipient)
       QC-93-10 Antibody screen or identification (donor/unit or recipient)
       QC-93-11 Antigen screen
       QC-93-12 Syphilis
       QC-93-13 All viral markers
       QC-93-14 Compatibility
       QC-93-15 HIV/HCV Nucleic Acid Test (NAT)
QC-94-** Inappropriate release of:
       QC-94-01 Other
       QC-94-02 Outdated product
       QC-94-03 Autologous unit not meeting homologous criteria
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QC-91-49 Incarcerated

QC-94-04 Product with unacceptable, undocumented, or incomplete product QC

QC-94-05 Product in which specification other than QC not met

QC-94-06 Product in which instrument QC or validation was unacceptable, incomplete, or not documented

QC-94-08 Product prior to resolution of discrepancy

QC-94-09 Product associated with product that contained clots or hemolysis

QC-94-12 Product identified as unsuitable due to a collection deviation or unexpected event

QC-94-13 Product identified as unsuitable due to a component preparation deviation or unexpected event

QC-94-14 Product identified as unsuitable due to a labeling deviation or unexpected event

QC-94-15 Product identified as unsuitable due to a donor screening deviation or unexpected event

QC-94-16 Product identified as unsuitable due to a donor deferral deviation or unexpected event

QC-96-** Shipping and storage

QC-96-01 Other

QC-96-02 Shipped at incorrect temperature

QC-96-03 Stored at incorrect temperature

QC-96-04 No documentation that product was shipped at appropriate temperature

QC-96-05 Temperature not recorded upon receipt, product redistributed

QC-96-06 Shipment exceeded time allowed for shipping, product redistributed

QC-96-07 Product not packaged for shipment in accordance with specifications

QC-97-** Distribution procedures not performed in accordance with blood bank transfusion service's specifications

OC-97-01 Other

QC-97-02 Product not irradiated as required

QC-97-03 Product issued to wrong patient

QC-97-04 Improper product selected for patient

QC-97-05 Improper ABO or Rh type selected for patient

QC-97-06 Product not leukoreduced as required

QC-97-07 Product released prior to obtaining current sample for ABO, Rh, antibody screen and/or compatibility testing

QC-97-08 Product not CMV negative as required

QC-97-10 Filter not issued with product or incorrect filter issued

QC-97-11 Product not irradiated and leukoreduced as required

QC-97-12 Product not irradiated and CMV negative as required

QC-97-13 Procedure for issuing product not performed or documented in accordance with specifications

QC-97-14 ABO and/or Rh retype of unit not performed or performed incorrectly

QC-97-15 Visual inspection not performed or documented

QC-97-16 Product inspected and accepted upon receipt from blood center, subsequently discovered to be hemolyzed

MI-**-** MISCELLANEOUS

MI-00-** Miscellaneous MI-00-01 Other

MI-01-** Donor implicated in transfusion associated disease

MI-01-01 Other MI-01-02 HIV MI-01-03 Hepatitis

MI-02-** Lookback; subsequent unit tested confirmed positive for:

MI-02-01 Other MI-02-02 HIV MI-02-04 HCV

??-??-?? DO NOT KNOW

Non-Blood BPD Codes

The following list of Biolo gical Product Deviation (BPD) Codes should be used as a tool for assigning a specific code to a reportable event when submitting the report to FDA. This list should not be used solely for the purpose of determining if an event is reportable. You should have a process consistent with the draft guidance document, "Biological Product Deviation Reporting for Manufacturers of Biological Products Other than Blood and Blood Components," to determine which events are reportable. The list includes deviations from regulations, standards, and standard operating procedures (SOPs) that may affect the safety, purity, or potency of a product. The list of codes was developed based on the error and accident reports previously submitted to FDA. These events may not apply to all establishments because they include deviations and unexpected events related to SOPs implemented at individual establishments and may not be an industry standard or a procedure at your facility. The use of BPD Codes will assist the FDA in analyzing the data submitted and streamline the trend analysis process.

IM - Incoming Material Specifications

PC - Process Controls

TE - Testing

LA - Labeling

PS - Product Specifications

QC - Quality Control and Distribution

MI - Miscellaneous

IM--*** INCOMING MATERIAL SPECIFICATIONS

IM-10-** Miscellaneous

IM-10-01 Other

IM-12-** Container

IM-12-01 Specifications not met

IM-12-02 Defective

IM-13-** Closures

IM-13-01 Specifications not met

IM-13-02 Defective

IM-14-** Source or raw material does not meet specifications or otherwise found to be unsuitable

IM-14-01 Other

IM-14-02 Contains precipitate

IM-14-03 Contaminated with microorganism

IM-14-04 Contaminated with mold

IM-14-05 Impurities exceed specification

IM-14-06 Testing deviation

IM-14-07 Stored or shipped at incorrect temperature or lack of controlled temperature

PC-**-** PROCESS CONTROLS

PC-20-** Miscellaneous

PC-20-01 Other

PC-21-** Manufacturing or processing performed using incorrect parameters

PC-21-01 Other

PC-21-02 Incorrect temperature

PC-21-03 Filling not performed according to specifications

PC-21-04 Aseptic processing procedures not performed according to specifications

PC-22-** Process/Procedure

PC-22-01 Other

PC-22-02 Interruption of process

PC-22-03 Environmental monitoring excursions

PC-22-04 Equipment not qualified

PC-22-05 Sanitization not performed or performed incorrectly

PC-22-06 Failed media fill

PC-23-** Process Water - specification not met

PC-23-01 Other

PC-23-02 Water for injection

PC-23-03 Purified water

PC-24-** Bulk material does not meet specifications or otherwise found to be unsuitable

PC-24-01 Other

PC-24-02 Contains precipitate

PC-24-03 Contaminated with microorganism

PC-24-04 Contaminated with mold

PC-24-05 Impurities exceed specification

PC-24-06 Stored at incorrect temperature

PC-24-07 Stored for an excessive hold time

TE-**-** TESTING

TE-30-** Miscellaneous

TE-30-01 Other

TE-31-** Safety

TE-31-01 Performed incorrectly

TE-31-02 Not performed or not documented

TE-32-** Purity

TE-32-01 Performed incorrectly

TE-32-02 Not performed or not documented

TE-33-** Potency

TE-33-01 Performed incorrectly

TE-33-02 Not performed or not documented

TE-34-** Sterility

TE-34-01 Performed incorrectly

TE-34-02 Not performed or not documented

TE-35-** Identity

TE-35-01 Performed incorrectly TE-35-02 Not performed or not documented

TE-36-** Stability

TE-36-01 Performed incorrectly TE-36-02 Not performed or not documented

LA-**-** LABELING

LA-40-** Miscellaneous LA-40-01 Other

LA-41-** Package insert

LA-41-01 Incorrect LA-41-02 Missing LA-41-03 Not current or approved

LA-42-** Product label

LA-42-01 Incorrect LA-42-02 Missing

LA-43-** Carton label

LA-43-01 Incorrect LA-43-02 Missing

LA-44-** Expiration date

LA-44-01 Extended LA-44-02 Missing

LA-45-** Lot number

LA-45-01 Incorrect LA-45-02 Missing

LA-46-** Storage temperature

LA-46-01 Incorrect LA-46-02 Missing

LA-47-** Administration route

LA-47-01 Incorrect LA-47-02 Missing

LA-48-** Concentration or volume

LA-48-01 Incorrect LA-48-02 Missing

PS--**** PRODUCT SPECIFICATIONS

PS-50-** Miscellaneous PS-50-01 Other

PS-51-** Product specification not met

PS-51-01 Other

PS-51-02 Contains precipitate

PS-51-03 Contaminated with microorganism

PS-51-04 Contaminated with mold

PS-51-05 Impurity levels

PS-51-06 Moisture

PS-51-07 Preservative content

PS-51-08 Potency

PS-51-09 Appearance

PS-51-10 Fill volume

PS-52-**Component packaged with final product did not meet specifications

PS-52-01 Other

PS-52-02 Contains precipitate

PS-52-03 Contaminated with microorganism

PS-52-04 Contaminated with mold

PS-52-05 Fill volume

PS-53-** Stability testing failed

PS-53-01 Other

PS-53-02 Potency

PS-53-03 Preservative

PS-53-04 Container closure integrity

PS-53-05 Chemical analysis/purity

PS-54-** Administration set (packaged with product) incorrect or incomplete

PS-54-01 Other

PS-54-02 Incorrect or missing label

PS-54-03 Defective PS-54-04 Expired

QC-**-** QUALITY CONTROL AND DISTRIBUTION

QC-60-** Miscellaneous QC-60-01 Other

QC-61-** Product distributed inappropriately

QC-61-01 Other

QC-61-02 Product distributed prior to completion of required testing

QC-61-03 Product distributed prior to CBER approval of a PAS

QC-61-04 Product distributed less than 30 days after submission of CBE-30

QC-61-05 Product distributed prior to validation of process

QC-61-06 Outdated product distributed

QC-62-** Shipping and storage

QC-62-01 Other

QC-62-02 Product shipped at incorrect temperature

QC-62-03 Product stored at incorrect temperature

QC-63-** Product identified as unacceptable, and not quarantined

QC-63-01 Other

MI-**-** MISCELLANEOUS

MI-70-** Miscellaneous MI-70-01 Other

??-??-?? DO NOT KNOW

Last Updated: 9/16/2002